equal the system must invalidate the order.

- (6) The system must check the Certificate Revocation List automatically and invalidate any order with a certificate listed on the Certificate Revocation List.
- (7) The system must check the validity of the certificate and the Certification Authority certificate and invalidate any order that fails these validity checks.
- (8) The system must have a time system that is within five minutes of the official National Institute of Standards and Technology time source.
- (9) The system must check the substances ordered against the schedules that the registrant is allowed to order and invalidate any order that includes substances the registrant is not allowed to order.
- (10) The system must ensure that an invalid finding cannot be bypassed or ignored and the order filled.
- (11) The system must archive the order and associate with it the digital certificate received with the order.
- (12) If a registrant sends reports on orders to DEA, the system must create a report in the format DEA specifies, as provided in §1305.29 of this chapter.
- (d) For systems used to process CSOS orders, the system developer or vendor must have an initial independent third-party audit of the system and an additional independent third-party audit whenever the signing or verifying functionality is changed to determine whether it correctly performs the functions listed under paragraphs (b) and (c) of this section. The system developer must retain the most recent audit results and retain the results of any other audits of the software completed within the previous two years.

### §1311.60 Recordkeeping.

- (a) A supplier and purchaser must maintain records of CSOS electronic orders and any linked records for two years. Records may be maintained electronically. Records regarding controlled substances that are maintained electronically must be readily retrievable from all other records.
- (b) Electronic records must be easily readable or easily rendered into a format that a person can read. They must

be made available to the Administration upon request.

(c) CSOS certificate holders must maintain a copy of the subscriber agreement that the Certification Authority provides for the life of the certificate.

# PART 1312—IMPORTATION AND EXPORTATION OF CONTROLLED SUBSTANCES

Sec.

1312.01 Scope of part 1312.

1312.02 Definitions.

IMPORTATION OF CONTROLLED SUBSTANCES

1312.11 Requirement of authorization to import.

1312.12 Application for import permit.

1312.13 Issuance of import permit.

1312.14 Distribution of copies of import permit.

1312.15 Shipments in greater or less amount than authorized.

1312.16 Cancellation of permit; expiration date.

1312.17 Special report from importers.

1312.18 Contents of import declaration.

1312.19 Distribution of import declaration.

EXPORTATION OF CONTROLLED SUBSTANCES

1312.21 Requirement of authorization to export.

1312.22 Application for export permit.

1312.23 Issuance of export permit.
1312.24 Distribution of copies of export permit.

1312.25 Expiration date.

1312.26 Records required of exporter.

1312.27 Contents of special controlled substances invoice.

1312.28 Distribution of special controlled substances invoice.

1312.29 Domestic release prohibited.

1312.30 Schedule III, IV, and V non-narcotic controlled substances requiring an import and export permit.

TRANSSHIPMENT AND IN-TRANSIT SHIPMENT OF CONTROLLED SUBSTANCES

1312.31 Schedule I: Application for prior written approval.

1312.32 Schedules II, III, IV: Advance notice.

#### HEARINGS

1312.41 Hearings generally.

1312.42 Purpose of hearing.

1312.43 Waiver or modification of rules.

1312.44 Request for hearing or appearance; waiver.

1312.45 Burden of proof.

1312.46 Time and place of hearing.

1312.47 Final order.

#### § 1312.01

AUTHORITY: 21 U.S.C. 952, 953, 954, 957, 958

SOURCE: 36 FR 7815, Apr. 24, 1971, unless otherwise noted. Redesignated at 38 FR 26609, Sept. 24, 1973.

#### §1312.01 Scope of part 1312.

Procedures governing the importation, exportation, transshipment and intransit shipment of controlled substances pursuant to section 1002, 1003, and 1004 of the Act (21 U.S.C. 952, 953, and 954) are governed generally by those sections and specifically by the sections of this part.

#### § 1312.02 Definitions.

Any term contained in this part shall have the definition set forth in section 102 of the Act (21 U.S.C. 802) or part 1300 of this chapter.

[62 FR 13969, Mar. 24, 1997]

IMPORTATION OF CONTROLLED SUBSTANCES

## § 1312.11 Requirement of authorization to import.

(a) No person shall import or cause to be imported any controlled substance listed in Schedule I or II or any narcotic controlled substance listed in Schedule III, IV or V or any non-narcotic controlled substance in Schedule III which the Administrator has specifically designated by regulation in §1312.30 of this part or any non-narcotic controlled substance in Schedule IV or V which is also listed in Schedule I or II of the Convention on Psychotropic Substances unless and until such person is properly registered under the Act (or exempt from registration) and the Administrator has issued him a permit to do so pursuant to §1312.13 of this part.

- (b) No person shall import or cause to be imported any non-narcotic controlled substance listed in Schedule III, IV or V, excluding those described in paragraph (a) of this section, unless and until such person is properly registered under the Act (or exempt from registration) and has filed an import declaration to do so with the Administrator, pursuant to § 1312.18 of this part.
- (c) When an import permit or declaration is required, a separate permit or declaration must be obtained for

each consignment of controlled substances to be imported.

[36 FR 7815, Apr. 24, 1971, as amended at 37 FR 15923, Aug. 8, 1972. Redesignated at 38 FR 26609, Sept. 24, 1973, and amended at 52 FR 17289, May 7, 1987]

## § 1312.12 Application for import permit.

(a) An application for a permit to import controlled substances shall be made on DEA Form 357. DEA Form 357 may be obtained from, and shall be filed with, the Drug Enforcement Administration, Drug Operations Section, Washington, DC 20537. Each application shall show the date of execution; the registration number of the importer: a detailed description of each controlled substance to be imported including the drug name, dosage form, National Drug Code (NDC) number, the Administration Controlled Substance Code Number as set forth in part 1308 of this chapter, the number and size of packages or containers, the name and quantity of the controlled substance contained in any finished dosage units, and the net quantity of any controlled substance (expressed in anhydrous acid, base or alkaloid) given in kilograms or parts thereof. The application shall also include the following:

- (1) The name, address, and business of the consignor, if known at the time application is submitted, but if unknown at that time, the fact should be indicated and the name and address afterwards furnished to the Administrator as soon as ascertained by the importer:
- (2) The foreign port of exportation (i.e., the place where the article will begin its journey of exportation to the United States);
- (3) The port of entry into the United States:
- (4) The latest date said shipment will leave said foreign port;
- (5) The stock on hand of the controlled substance desired to be imported;
- (6) The name of the importing carrier or vessel (if known, or if unknown it should be stated whether shipment will be made by express, freight, or otherwise, imports of controlled substances in Schedules I or II and narcotic drugs